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# Supplementary data

### **Supplementary text**

#### Cloning of SmAOS2

Total RNA of *S. moellendorffii* (ca. 120 mg) was extracted using the conventional phenol-chloroform method. Reverse transcription (M-MLV Reverse Transcriptase, Invitrogen, Carlsbad, CA USA) was performed according to the manufacturer's instructions to generate cDNA. The primer set of *SmAOS2*, SmAOS2-F and SmAOS2-R, was used to amplify the open reading frame (ORF) of *SmAOS2*. To afford the ORF of *SmAOS2*, the PCR reaction was performed in 50 μl per tube, containing 10 μl of dNTP mixture (2.0 mM each), 1 μl each from forward and reverse primer (5 μM), 0.5 μl of cDNA, 25 μl of PCR buffer (2 ×), 0.5 μl of KOD FX DNA polymerase (Toyobo, Japan) and 12 μl of Milli-Q water under following conditions; pre-denaturation at 94°C for 2 min followed by 40 cycles of denaturing at 98°C for 10 s, annealing at 56°C for 30 s, and extension at 68°C for 1 min, and then a final extension at 68°C for 10 min. The obtained PCR product was purified and inserted into the pBlueScript SK II (+) vector (Stratagene, USA), which was digested with *EcoRV* to obtain the plasmid of pSK-SmAOS2. DNA sequencing was performed to confirm that the target gene was successfully inserted into the plasmid. The primers used in this study are listed in Supplementary Table S2.

#### Synthesis of a recombinant SmAOS2 in E. coli

The primer set for overexpression of SmAOS2, SmAOS2NdeI-F with NdeI site and SmAOS2XhoI-R with XhoI site, in E. coli was designed according to a SmAOS2 sequence. PCR was performed in 15 µl of a reaction mixture containing 1.5 µl of dNTP mixture (2.0 mM each dNTP), 0.9 µl of 25 mM MgSO<sub>4</sub>, 1.5 µl of KOD plus neo buffer (10 ×), 0.9 µl of each primer (SmAOS2NdeI-F and SmAOS2XhoI-R, 5 µM), 1 µl of pSK-SmAOS2 (200 ng/µl), 0.3 μl of KOD plus neo DNA polymerase (Toyobo, Japan), and 8 μl of Milli-Q water. PCR was conducted with the following conditions: 2 min at 94°C for pre-denaturation; 30 cycles of 98°C for 10 s, 58°C for 30 s, and 68°C for 90 s and then final extension at 68°C for 10 min. The PCR product then was purified and ligated into pET23a vector (Merck, USA) using Ligation Mix (Takara, Japan) according to the standard procedure. The constructed plasmid, pET23a-SmAOS2, was transformed in E.coli BL21 (DE3) according to standard procedure and was subsequently grown in LB agar medium supplemented with 100 µg/ml of ampicillin. A single colony was inoculated into 10 ml of LB medium containing 100 µg/ml of ampicillin and then incubated at 37°C for overnight. A 10 ml aliquot was transferred into 1 l of LB medium containing 100 µg/ml of ampicillin and then incubated at 37°C until the OD<sub>600</sub> reached 0.4; the sample was finally induced by 1 mM of IPTG for overnight at 25°C. The bacterial cells were collected by centrifugation at  $5,000 \times g$  for 20 min and lysed by ultrasonication in 50 mM phosphate buffer pH 7.4, 0.3 M NaCl, and 20 mM imidazole. Cell debris was removed by centrifugation at  $15,000 \times g$  for 15 min; then, the supernatant was subjected to Ni-NTA agarose column chromatography (2 ml, GE Healthcare, USA). SmAOS2, fused with a His-tag, was eluted with 50 mM phosphate buffer (pH 7.8) containing 100 mM imidazole and 0.3 M NaCl. The resulting recombinant protein solution was dialyzed with 50 mM Tris-HCl (pH 8.0) containing 20 mM NaCl. The purity was checked by SDS-PAGE (Supplementary Fig. S3). The dialyzed product was concentrated using Amicon Ultra-15 Centrifugal Filter Device 10K (Merck Millipore Ltd., Germany) and then used for evaluation of enzymatic activity. The primers used in this study are listed in Supplementary Table S2.

#### Cloning of SmAOC1

Total RNA and cDNA of *S. moellendorffii* were obtained according the method described above. Full length cDNA of *SmAOC1* (Sm\_91887) was obtained with 5'-Full RACE Core Set according to the manufacturer's instructions. Subsequently, the primer set of *SmAOC1*, SmAOC1-F and SmAOC1-R was used to amplify the open reading frame (ORF) of *SmAOC1*. The PCR reaction was performed in 50 μl per tube, containing 10 μl of dNTP mixture (2.0 mM each), 1 μl each from forward and reverse primer (5 μM), 0.5 μl of cDNA, 25 μl of PCR buffer (2 ×), 0.5 μl of KOD FX DNA polymerase (Toyobo, Japan) and 12 μl of Milli-Q water under following conditions: predenaturation at 94°C for 2 min followed by 40 cycles of denaturing at 98°C for 10 s, annealing at 55°C for 30 s, and extension at 68°C for 1 min, and then a final extension at 68°C for 10 min. The obtained PCR product was purified and inserted into the pBlueScript SK II (+) vector (Stratagene, USA), which was digested with *EcoRV* to obtain the plasmid of pSK-SmAOC1. DNA sequencing was performed to confirm that the target gene was successfully inserted into the plasmid. The primers used in this study are listed in Supplementary Table S2.

#### Synthesis of a recombinant SmAOC1 in E. coli

ChloroP v 1.1 and TargetP predicted that the fifty amino acids at the N-terminus of SmAOC1 was a chloroplast transit peptide. The primer set for overexpression of SmAOC1, SmAOC1SphI-F with SphI site and SmAOC1SalI-R with Sall site, in E. coli was designed according to a SmAOC1 sequence, which had 150 nucleotides deleted from the codon for the first methionine. PCR was performed in 15 µl of a reaction mixture containing 1.5 µl of dNTP mixture (2.0 mM each dNTP), 0.9 µl of 25 mM MgSO<sub>4</sub>, 1.5 µl KOD plus neo buffer (10 ×), 0.9 µl of each primer (SmAOC1SphI-F and SmAOC1SalI-R, 5 µM), 1 µl of pSK-SmAOC1 (200 ng/µl), 0.3 µl of KOD plus neo DNA polymerase (Toyobo, Japan), and 8 µl of Milli-Q water. PCR was conducted with the following conditions: 2 min at 94°C for pre-denaturation; 30 cycles of 98°C for 10 s, 60°C for 30 s, and 68°C for 90 s and then final extension at 68°C for 10 min. The PCR product then was purified and ligated into pQE30 vector (Qiagen, USA) using T4 DNA ligase (Takara, Japan) according to the standard procedure. The constructed plasmid, pQE30-SmAOC1, was transformed in E.coli M15 according to standard procedure and was subsequently grown in LB agar medium supplemented with 100 µg/ml of ampicillin. A single colony was inoculated into 10 ml of LB medium containing 100 µg/ml of ampicillin and then incubated at 37°C for overnight. A 10 ml aliquot was transferred into 1 l of LB medium containing 100 µg/ml of ampicillin and then incubated at 25°C until the OD<sub>600</sub> reached 0.5; the sample was finally induced by 0.2 mM of IPTG for 4 h at 25°C. The bacterial cells were collected by centrifugation at  $7,000 \times g$ for 20 min and lysed by ultrasonication in 50 mM phosphate buffer pH 7.8, 0.3 M NaCl, and 20 mM imidazole. Cell debris was removed by centrifugation at 15,000 × g for 15 min; then, the supernatant was subjected to Ni-NTA agarose column chromatography (2 ml, GE Healthcare, USA). SmAOC1, fused with a His-tag, was eluted with 50 mM phosphate buffer (pH 7.8) containing 200 mM imidazole and 0.3 M NaCl. The resulting recombinant protein solution was dialyzed with 50 mM Tris-HCl (pH 8.0) containing 20 mM NaCl. The purity was checked by SDS-PAGE (Supplementary Fig. S3). The dialyzed product was concentrated using Amicon Ultra-15 Centrifugal Filter Device 10K (Merck Millipore Ltd., Germany) and then used for evaluation of enzymatic activity. The primers used in this study are listed in Supplementary Table S2.

#### Subcellular localization of SmAOC1

To analyze the localization of SmAOC1, PCR was performed using primers, SmAOC1NdeI-F and SmAOC1EcoRV-R, with *NdeI* and *EcoRV* restriction sites, respectively. The PCR reaction mixture contained 1.5  $\mu$ l of dNTP mixture (2.0 mM each dNTP), 0.9  $\mu$ l of 25 mM MgSO<sub>4</sub>, 1.5  $\mu$ l of KOD plus neo buffer (10 ×), 0.9  $\mu$ l of each primer (5  $\mu$ M), 1  $\mu$ l of pSK-SmAOC1 (200 ng/ $\mu$ l), 0.3  $\mu$ l of KOD plus neo DNA polymerase (Toyobo, Japan), and an adjusted Milli-Q water level up to a volume of 15  $\mu$ 1. PCR was conducted for 2 min at 94°C for pre-

denaturation; 30 cycles of 98°C for 10 s, 60°C for 30 s, and 68°C for 90 s; final extension was performed at 68°C for 10 min. The PCR product was ligated into the pENTR4 vector using GeneArt Seamless Cloning and Assembly Kit (Invitrogen, USA). The pENTR4-SmAOC1 entry clone was introduced into the pUGWnew5 destination vector by Gateway LR Clonase Mix (Invitrogen, USA) to generate the plasmid encoding 35S::SmAOC1-GFP (pUGWnew5-SmAOC1). The constructed plasmid of 35S::SmAOC1-GFP was transformed into protoplasts of *P. patens*, which were grown in BCDATG medium for 3 days under white fluorescent light using the PEG-mediated transformation technique. Localization of the SmAOC1-GFP fusion protein was observed under a TCS-SP5 confocal laser scanning microscope (Leica, Germany) after 3 days incubation. Images were observed at an excitation of 488 nm and emission of 530 nm for detecting the GFP signal as well as emission over 655 nm for detecting auto-fluorescence from the chloroplasts. The primers used in this study are listed in Supplementary Table S2.

#### Cloning of SmOPR1

Total RNA and cDNA of *S. moellendorffii* were obtained according the method described above. The primer set of *SmOPR1*, SmOPR1-F and SmOPR1-R, was used to amplify the open reading frame (ORF) of *SmOPR1*. To afford the ORF of *SmOPR1*, the PCR reaction was performed in 50 μl per tube, containing 10 μl of dNTP mixture (2.0 mM each), 1 μl each from forward and reverse primer (5 μM), 0.5 μl of cDNA, 25 μl of PCR buffer (2 ×), 0.5 μl of KOD FX DNA polymerase (Toyobo, Japan) and 12 μl of Milli-Q water under following conditions; pre-denaturation at 94°C for 2 min followed by 40 cycles of denaturing at 98°C for 10 s, annealing at 60°C for 30 s, and extension at 68°C for 1 min, and then a final extension at 68°C for 10 min. The obtained PCR product was purified and inserted into the pBlueScript SK II (+) vector (Stratagene, USA), which was digested with *EcoRV* to obtain the plasmid of pSK-SmOPR1. DNA sequencing was performed to confirm that the target gene was successfully inserted into the plasmid. The primers used in this study are listed in Supplementary Table S2.

#### Synthesis of a recombinant SmOPR1 in E. coli

The primer set for overexpression of SmOPR1, SmOPR1EcoRI-F with EcoRI site and SmOPR1NotI-R with NotI site, in E. coli was designed according to a SmOPR1 sequence. PCR was performed in 15 µl of a reaction mixture containing 1.5 µl of dNTP mixture (2.0 mM each dNTP), 0.9 µl of 25 mM MgSO<sub>4</sub>, 1.5 µl of KOD plus neo buffer (10 ×), 0.9 μl of each primer (SmOPR1EcoRI-F and SmOPR1NotI-R, 5 μM), 1 μl of pSK-SmOPR1 (200 ng/ μl), 0.3 μl of KOD plus neo DNA polymerase (Toyobo, Japan), and 8 μl of Milli-Q water. PCR was conducted with the following conditions: 2 min at 94°C for pre-denaturation; 30 cycles of 98°C for 10 s, 60°C for 30 s, and 68°C for 90 s and then final extension at 68°C for 10 min. The PCR product then was purified and ligated into pET23a vector (Merck, USA) using Ligation Mix (Takara, Japan) according to the manufacturer's instruction. The constructed plasmid, pET23a-SmOPR1, was transformed in E.coli BL21 (DE3) according to standard procedure and was subsequently grown in LB agar medium supplemented with 100 µg/ml of ampicillin. A single colony was inoculated into 10 ml of LB medium containing 100 µg/ml of ampicillin and then incubated at 37°C for overnight. A 10 ml aliquot was transferred into 1 l of LB medium containing 100 µg/ml of ampicillin and then incubated at 25°C until the OD<sub>600</sub> reached 0.4; the sample was finally induced by 1 mM of IPTG for overnight at 18°C. The bacterial cells were collected by centrifugation at  $7,000 \times g$  for 20 min and lysed by ultrasonication in 50 mM phosphate buffer pH 7.4, 0.3 M NaCl, and 20 mM imidazole. Cell debris was removed by centrifugation at  $15,000 \times g$  for 15 min; then, the supernatant was subjected to Ni-NTA agarose column chromatography (2 ml, GE Healthcare, USA). SmOPR1, fused with a His-tag, was eluted with 50 mM phosphate buffer (pH 7.4) containing 200 mM imidazole and 0.3 M NaCl. The resulting recombinant protein solution was dialyzed with 50 mM Tris-HCl (pH 7.8) containing 20 mM NaCl. The purity was checked by SDS-PAGE (Supplementary Fig. S3). The dialyzed product was concentrated using

Amicon Ultra-0.5mL Centrifugal Filter Units 30K and 50K (Merck Millipore Ltd., Germany) and then used for evaluation of enzymatic activity. The primers used in this study are listed in Supplementary Table S2.

#### Cloning of SmOPR5

Total RNA and cDNA of *S. moellendorfii* were obtained according the method described above. The primer set of *SmOPR5*, SmOPR5-F and SmOPR5-R, was used to amplify the open reading frame (ORF) of *SmOPR5*. To afford the ORF of *SmOPR5*, the PCR reaction was performed in 50 μl per tube, containing 10 μl of dNTP mixture (2.0 mM each), 1 μl each from forward and reverse primer (5 μM), 0.5 μl of cDNA, 25 μl of PCR buffer (2 ×), 0.5 μl of KOD FX DNA polymerase (Toyobo, Japan) and 12 μl of Milli-Q water under following conditions; pre-denaturation at 94°C for 2 min followed by 40 cycles of denaturing at 98°C for 10 s, annealing at 60°C for 30 s, and extension at 68°C for 1 min, and then a final extension at 68°C for 10 min. The obtained PCR product was purified and inserted into the pBlueScript SK II (+) vector (Stratagene, USA), which was digested with *EcoRV* to obtain the plasmid of pSK-SmOPR5. DNA sequencing was performed to confirm that the target gene was successfully inserted into the plasmid. The primers used in this study are listed in Supplementary Table S2.

#### Synthesis of a recombinant SmOPR5 in E. coli

The primer set for overexpression of SmOPR5, SmOPR5BamHI-F with BamHI site and SmOPR5XhoI-R with XhoI site, in E. coli was designed according to a SmOPR5 sequence. PCR was performed in 15 µl of a reaction mixture containing 1.5 µl of dNTP mixture (2.0 mM each dNTP), 0.9 µl of 25 mM MgSO<sub>4</sub>, 1.5 µl of KOD plus neo buffer (10 ×), 0.9 µl of each primer (SmOPR5BamHI-F and SmOPR5XhoI-R, 5 µM), 1 µl of pSK-SmOPR5 (200 ng/ μl), 0.3 μl of KOD plus neo DNA polymerase (Toyobo, Japan), and 8 μl of Milli-Q water. PCR was conducted with the following conditions: 2 min at 94°C for pre-denaturation; 30 cycles of 98°C for 10 s, 60°C for 30 s, and 68°C for 90 s and then final extension at 68°C for 10 min. The PCR product then was purified and ligated into pET23a vector (Merck, USA) using In-Fusion HD Cloning Kit (Takara, Japan) according to the manufacturer's instruction. The constructed plasmid, pET23a-SmOPR5, was transformed in E.coli BL21 (DE3) according to standard procedure and was subsequently grown in LB agar medium supplemented with 100 µg/ml of ampicillin. A single colony was inoculated into 10 ml of LB medium containing 100 µg/ml of ampicillin and then incubated at 37°C for overnight. A 10 ml aliquot was transferred into 1 l of LB medium containing 100 μg/ml of ampicillin and then incubated at 25°C until the OD<sub>600</sub> reached 0.4; the sample was finally induced by 1 mM of IPTG for overnight at 18°C. The bacterial cells were collected by centrifugation at 15,000 × g for 20 min and lysed by ultrasonication in 50 mM phosphate buffer pH 7.4, 0.3 M NaCl, and 20 mM imidazole. Cell debris was removed by centrifugation at 15,000 x g for 15 min; then, the supernatant was subjected to Ni-NTA agarose column chromatography (2 ml, GE Healthcare, USA). SmOPR5, fused with a His-tag, was eluted with 50 mM phosphate buffer (pH 7.4) containing 200 mM imidazole and 0.3 M NaCl. The resulting recombinant protein solution was dialyzed with 50 mM Tris-HCl (pH 7.8) containing 20 mM NaCl. The purity was checked by SDS-PAGE (Supplementary Fig. S3). The dialyzed product was concentrated using Amicon Ultra-0.5ml Centrifugal Filter Units 30K and 50K (Merck Millipore Ltd., Germany) and then used for evaluation of enzymatic activity. The primers used in this study are listed in Supplementary Table S2.

#### Cloning of SmJAR1

Total RNA and cDNA of *S. moellendorfii* were obtained according the method described above. The primer set of *SmJAR1*, SmJAR1-F and SmJAR1-R, was used to amplify the open reading frame (ORF) of *SmJAR1*. To afford the ORF of *SmJAR1*, the PCR reaction was performed in 50  $\mu$ l per tube, containing 10  $\mu$ l of dNTP mixture (2.0 mM each), 1  $\mu$ l each from forward and reverse primer (5  $\mu$ M), 0.5  $\mu$ l of cDNA, 25  $\mu$ l of PCR buffer (2  $\times$ ), 0.5  $\mu$ l of KOD

FX DNA polymerase (Toyobo, Japan) and 12 μl of Milli-Q water under following conditions; pre-denaturation at 94°C for 2 min followed by 40 cycles of denaturing at 98°C for 10 s, annealing at 60°C for 30 s, and extension at 68°C for 1 min, and then a final extension at 68°C for 10 min. The obtained PCR product was purified and inserted into the pBlueScript SK II (+) vector (Stratagene, USA), which was digested with *EcoRV* to obtain the plasmid of pSK-SmJAR1. DNA sequencing was performed to confirm that the target gene was successfully inserted into the plasmid. The primers used in this study are listed in Supplementary Table S2.

### Construction of E. coli transferred with an overexpression vector of SmJAR1

The primer set for overexpression of SmJAR1, SmJAR1BamHI-F with *BamHI* site and SmJAR1XhoI-R with *XhoI* site, in *E. coli* was designed according to a *SmJAR1* sequence. PCR was performed in 15 μl of a reaction mixture containing 1.5 μl of dNTP mixture (2.0 mM each dNTP), 0.9 μl of 25 mM MgSO<sub>4</sub>, 1.5 μl of KOD plus neo buffer (10 ×), 0.9 μl of each primer (SmJAR1BamHI-F and SmJAR1XhoI-R, 5 μM), 1 μl of pSK-SmJAR1 (200 ng/ μl), 0.3 μl of KOD plus neo DNA polymerase (Toyobo, Japan), and 8 μl of Milli-Q water. PCR was conducted with the following conditions: 2 min at 94°C for pre-denaturation; 30 cycles of 98°C for 10 s, 60°C for 30 s, and 68°C for 90 s and then final extension at 68°C for 10 min. The PCR product then was purified and ligated into pET23a vector (Merck, USA) using In-Fusion HD Cloning Kit (Takara, Japan) according to the manufacturer's instruction. The constructed plasmid, pET23a-SmJAR1, was transformed in *E.coli* BL21 (DE3) according to standard procedure. The resultant strain was used for an assay of JA-Ile synthetic activity. The primers used in this study are listed in Supplementary Table S2.

# **Supplementary tables**

Supplementary Table S1. Putative genes of AOS, AOC, OPR, and JAR1 in S. moellendorffii.

Designation	Locus name	Location	Query (GenBank accession no.)
SmAOS1	271334	scaffold_38	Arabidopsis AOS (AED94842)
SmAOS2	177201	scaffold_38	Arabidopsis AOS (AED94842)
SmAOS3	228572	scaffold_38	Arabidopsis AOS (AED94842)
SmAOC1	91887	scaffold_12	Arabidopsis AOC (CAC83764)
SmOPR1*	270843	scaffold_14	Arabidopsis OPR3 (AEC06000)
SmOPR5*	111662	scaffold_41	Arabidopsis OPR3 (AEC06000)
SmJAR1	110439	scaffold_40	Arabidopsis JAR1 (AEC10684)

<sup>\*</sup>The numbering of SmOPRs was previously described by Li et al. (2009).

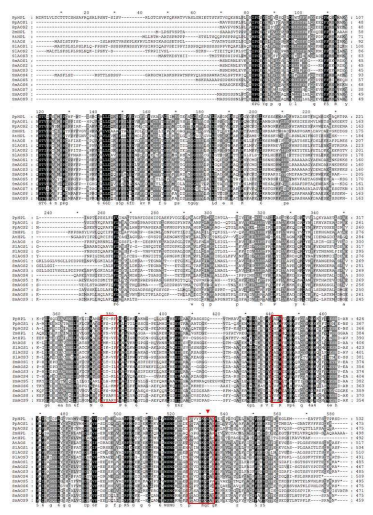
# Supplementary Table S2. Oligonucleotide primers.

Primer Name	Sequence 5' to 3'	Purpose	
SmAOC1-RT	GGAGCAACGGACTGC	5'-RACE PCR	
SmAOC1_A1	GAGATGTGGCCGTAGTCCC	5'-RACE PCR	
SmAOC1_S1	GGATACGTACATGGCGATCA	5'-RACE PCR	
SmAOC1_A2	AAGCTGTAGGTGGCCTCGTA	5'-RACE PCR	
SmAOC1_S2	CTTCTACTTGGAGGGCATCG	5'-RACE PCR	
SmAOC1-F	ATGGCAAGTTCCCTGGCG	ORF SmAOC1	
SmAOC1-R	TTAATCTGTGAAGTTTGGAGCAAC	ORF SmAOC1	
SmAOC1SphI-F	TAGCATGCTCAGCTGCCATTGTCCC	Hetero-overexpression of SmAOC1	
SmAOC1SalI-R	CGGTCGACTTAATCTGTGAAGTTTGGAG	Hetero-overexpression of SmAOC1	
SmAOC1NdeI-F	AAAAGCAGGCTCCACCATGGCAAGTTCCCT	Sub-cellular localization of SmAOC1 Sub-cellular localization of SmAOC1	
SmAOC1EcoRV-R	AAGCTGGGTCTAGATTTAATCTGTGAAGTT		
SmOPR1-F	ATGGATGCGCCCCAGGAGCA	ORF SmOPR1	
SmOPR1-R	AGCTATCGTTTTTCTTAATCTTCAAGGAAAGGA	ORF SmOPR1	
SmOPR1EcoRI-F	GAATTCATGGATGCGCCCCAGG	Hetero-overexpression SmOPR1 Hetero-overexpression SmOPR1	
SmOPR1NotI-R	GCGGCCGCATCTTCAAGGAAAGG		
SmOPR5-F	ATGGAAAGCTCATCAAATCCTCTGA	ORF SmOPR5	
SmOPR5-R	CTAAAGTTTGCTGGGGTGTTTCTT	ORF SmOPR5	
SmOPR5BamHI-F	ATGGGTCGCGGATCCATGGAAAGCTCATCA	Hetero-overexpression SmOPR5	
SmOPR5IXhoI-R	GTGGTGCTCGAGAAGTTTGCTGGGGTG	Hetero-overexpression SmOPR5	
SmJAR1-F	ATGCCAGGGATTCCATTGAT	ORF SmJAR1	
SmJAR1-R	CTACTCTCCTCACTCCCG	ORF SmJAR1	
SmJAR1BamHI-F	ATGGGTCGCGGATCCATGCCAGGGATTCCA	Hetero-overexpression SmJAR1	
SmJAR1IXhoI-R	GTGGTGGTGCTCGAGCTCTCTCCTCACTCC	Hetero-overexpression SmJAR1	

# Supplementary Table S2. Oligonucleotide primers (continued).

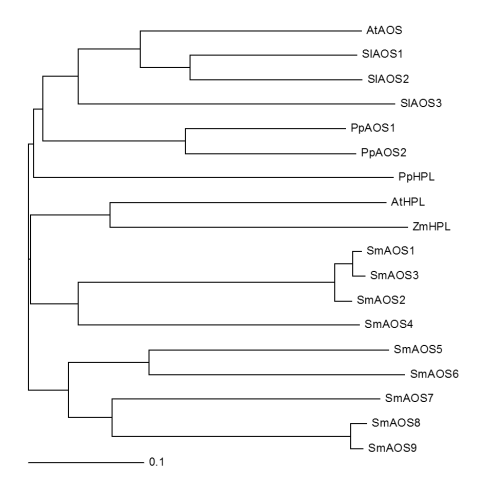
Primer Name	Sequence 5' to 3'	Purpose	
SmAOS1_F	CTTAGCAGCTCACACTTCAC	ORF SmAOS1	
SmAOS1_R	GACATACACACTAGTGCATTTC	ORF SmAOS1	
SmAOS2_F	CTTAGCAGCTCATACTTCACTTC	ORF SmAOS2	
SmAOS2_R	GACAACGAACACGGTGATTGTG	ORF SmAOS2	
SmAOS3_F	CTTAGCAGCTCAAACTTCAC	ORF SmAOS3	
SmAOS3_R	GCATTTCATTTCATGCTCGC	ORF SmAOS3	
SmAOS1EcoRI-F	CCGAATTCATGAGCAACGACAGGAACCT	Hetero-overexpression SmAOS1	
SmAOS1XhoI-R	TGCTCGAGTGCTCGCTTCTTGAGCTCGG	Hetero-overexpression SmAOS1	
SmAOS2NdeI-F	TACATATGAGCAACGACAGGAACCT	Hetero-overexpression SmAOS2	
SmAOS2XhoI-R	TGCTCGAGTGCTCGCTTCTTGAGCTCGG	Hetero-overexpression SmAOS2	
qSmAOC1-F	TTCCCGACCAAGCTCTTCTA	Expression analysis of SmAOC1	
qSmAOC1-R	AAGTTTGGAGCAACGGACTG	Expression analysis of SmAOC1	
qSmOPR5-F	AGCTGTGCATGACAAAGGTG	Expression analysis of SmOPR5	
qSmOPR5-R	GGACTGTCCATCTGGGAAGA	Expression analysis of SmOPR5	
qSmJAR1-F	AGTATACCGCCCATGCTGAC	Expression analysis of SmJAR1	
qSmJAR1-R	CCATGCAATCACAACACTCC	Expression analysis of SmJAR1	
qSmUbi1-F	ATACCATCGGCGATTTGAAG	Reference gene ubiquitin	
qSmUbi1-R	CGCTTACAAGGAAAGCACCT	Reference gene ubiquitin	
SmActin-F	ACTGGGACGACATGGAGAAG	Reference gene actin	
SmActin-R	CGCCTGAATAGCAACGTACA	Reference gene actin	

## **Supplementary figures**



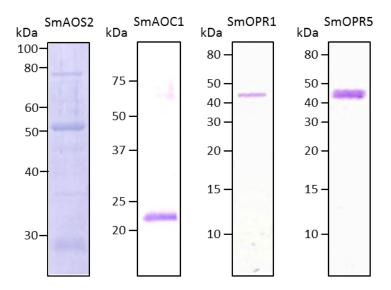
Supplementary Fig. S1. Amino acid sequence alignment of SmAOSs with the previously reported AOSs.

The Clustal Omega was used for the alignment. The conserved cysteine residue for heme-ligand binding is indicated with a red triangle. The I-helix GXXX(F/L), EXLR motif, and PXVXNKQCPG for heme-binding domain are in red box (Koeduka et al. 2015). The reported AOSs: PpAOS1 (Physcomitrella patens, CAC86919), PpAOS2 (Physcomitrella patens, XP\_001759629), SIAOS1 (Solanum lycopersicum, CAB88032), SIAOS2 (Solanum lycopersicum, AAF67141), SIAOS3 (Solanum lycopersicum, AAN76867), AtAOS (Arabidopsis thaliana, CAA63266), AtHPL (A. thaliana hydroperoxide lyase, AAC69871), ZmHPL(Zea mays hydroperoxide lyase, AAS47027), and PpHPL (P. patens hydroperoxide lyase, CAC86920). AOS homologues in Selaginella (SELMODRAFT 271334), moellendorffii: SmAOS1 SmAOS2 (SELMODRAFT 177201), SmAOS3 (SELMODRAFT 228572), SmAOS4 (SELMODRAFT 133317), SmAOS5 (SELMODRAFT 81998), SmAOS6 (SELMODRAFT\_177485), SmAOS7 (SELMODRAFT\_92382), SmAOS8 (SELMODRAFT\_98717), and SmAOS9 (SELMODRAFT\_41357).



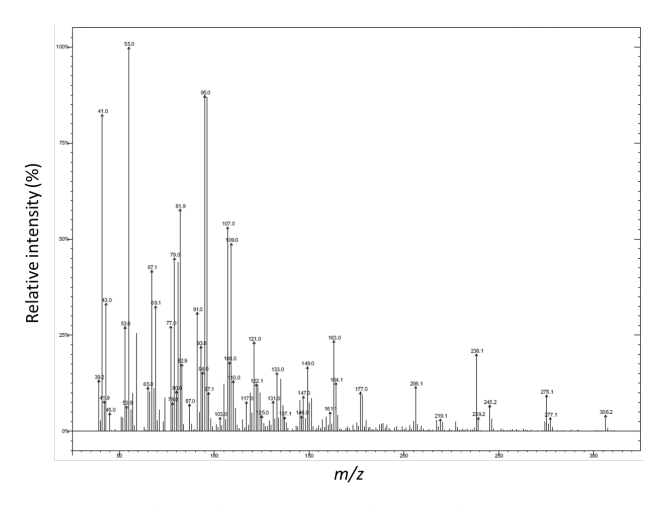
Supplementary Fig. S2. Phylogenetic tree of SmAOSs and previously reported AOSs.

Phylogenetic tree was constructed using TreeView X program based on amino acid sequence alignment. The bars represent evolutionary distance. The reliability of the tree measured by bootstrap analysis with 1,000 replicates. The reported AOSs: PpAOS1 (*Physcomitrella patens*, CAC86919), PpAOS2 (*Physcomitrella patens*, XP\_001759629), SIAOS1 (*Solanum lycopersicum*, CAB88032), SIAOS2 (*Solanum lycopersicum*, AAF67141), SIAOS3 (*Solanum lycopersicum*, AAN76867), AtAOS (*Arabidopsis thaliana*, CAA63266), AtHPL (*A. thaliana* hydroperoxide lyase, AAC69871), ZmHPL(*Zea mays* hydroperoxide lyase, AAS47027), and PpHPL (*P. patens* hydroperoxide lyase, CAC86920). AOS homologues in *Selaginella moellendorffii*: SmAOS1 (SELMODRAFT\_271334), SmAOS2 (SELMODRAFT\_177201), SmAOS3 (SELMODRAFT\_228572), SmAOS4 (SELMODRAFT\_133317), SmAOS5 (SELMODRAFT\_81998), SmAOS6 (SELMODRAFT\_177485), SmAOS7 (SELMODRAFT\_92382), SmAOS8 (SELMODRAFT\_98717), and SmAOS9 (SELMODRAFT\_41357).

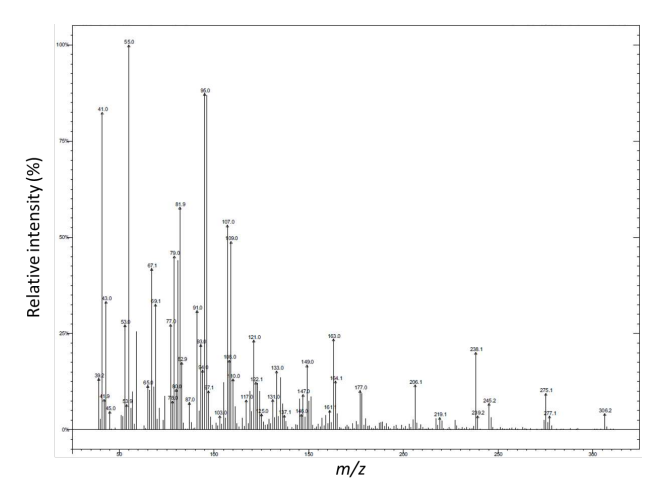


Supplementary Fig. S3. SDS-PAGE analysis of recombinant SmAOS2, SmAOC1, SmOPR1 and SmOPR5.

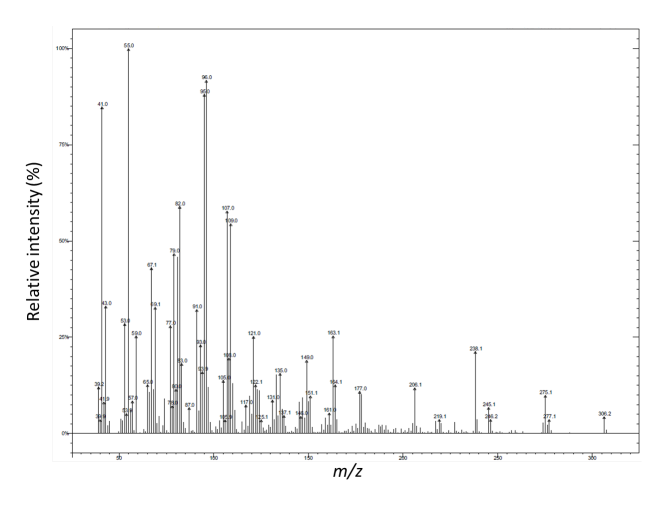
The recombinant proteins fused with His-tag were overexpressed in *E. coli* and were purified using Ni-NTA agarose column chromatography. SmAOC1 was analyzed by 15% SDS-PAGE. SmAOS1, SmOPR1, and SmOPR5 were analyzed by 10% SDS-PAGE, respectively. Proteins were stained by Coomassie Brilliant Blue (CBB).



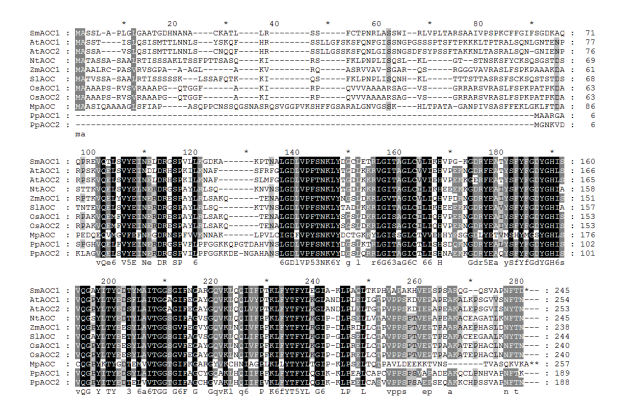
Supplementary Fig. S4. GC-MS spectrum of methyl ester of (+)-trans-OPDA.



Supplementary Fig. S5. GC-MS spectrum of methyl ester of (–)-*trans*-OPDA.

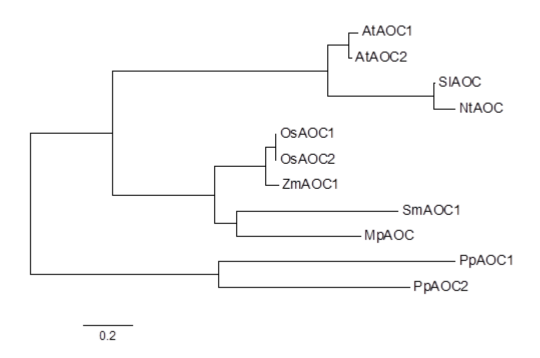


Supplementary Fig. S6. GC-MS spectrum of methyl ester of standard OPDA.



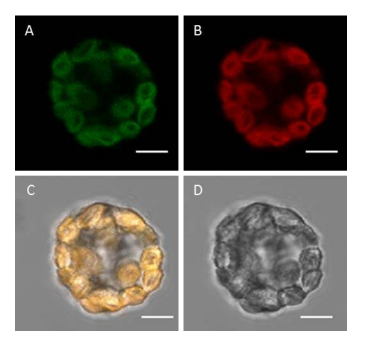
Supplementary Fig. S7. Amino acid sequence alignment of SmAOC1 with the previous reported AOCs.

Amino acid sequences were aligned using Clustal Omega. Identical and similar amino acids are highlighted in black and gray, respectively. The aligned sequences include SmAOC1 (*Selaginella moellendorffii*, Sm\_91887), PpAOC1 (*Physcomitrella patens*, CAD48752), PpAOC2 (*Physcomitrella patens*, CAD48753), MpAOC (*Marchantia polymorpha*, BAO93687), AtAOC1 (*Arabidopsis thaliana*, AEE77065.1), AtAOC2 (*Arabidopsis thaliana*, AEE77066.1), ZmAOC (*Zea mays*, NP\_001105245), OsAOC1 (*Oryza sativa*, ABV03555), OsAOC2 (*Oryza sativa*, ABV45432), SlAOC (*Solanum lycopersicum*, AAK62358), and NtAOC (*Nicotiana tabacum*, CAC83765).



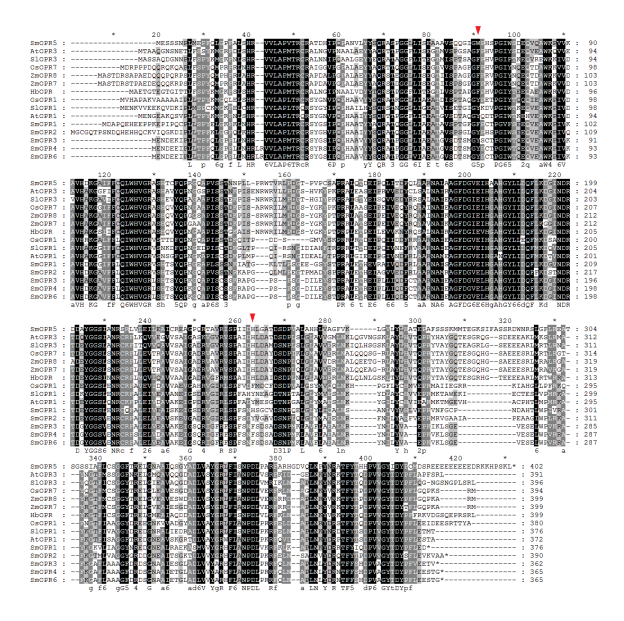
Supplementary Fig. S8. Phylogenetic tree of SmAOC1 and previously reported AOCs.

Phylogenetic tree was constructed using the neighbor-joining method with MEGA 5.2 program based on amino acid sequence alignment. The bars represent evolutionary distance. The reliability of the tree measured by bootstrap analysis with 1,000 replicates. The analysis was performed with following: SmAOC1 (*Selaginella moellendorffii*, Sm\_91887), PpAOC1 (*Physcomitrella patens*, CAD48752), PpAOC2 (*Physcomitrella patens*, CAD48753), MpAOC (*Marchantia polymorpha*, BAO93687), AtAOC1 (*Arabidopsis thaliana*, AEE77065.1), AtAOC2 (*Arabidopsis thaliana*, AEE77066.1), ZmAOC (*Zea mays*, NP\_001105245), OsAOC1 (*Oryza sativa*, ABV03555), OsAOC2 (*Oryza sativa*, ABV45432), SIAOC (*Solanum lycopersicum*, AAK62358), and NtAOC (*Nicotiana tabacum*, CAC83765).



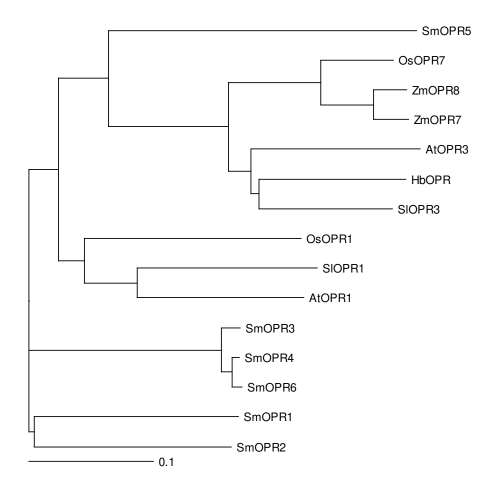
Supplementary Fig. S9. Expression of SmAOC1-GFP fusion protein in the chloroplast of *P. patens* protoplast.

The sub-cellular localization of SmAOC1 was analyzed by constructing the SmAOC1-GFP plasmid, which was introduced into the prepared protoplast of *P. patens* using PEG-mediated transformation method. Images were taken with a confocal laser-scanning microscope with an excitation of 488 nm and emission of 530 nm for detecting GFP and an emission above 655 nm for detecting autofluorescence from chlorophyll. (A) Green fluorescence of SmAOC1-GFP. (B) Red chlorophyll autofluorescence. (C) Merge of the green fluorescence and the red chlorophyll autofluorescence. (D) Bright field. Scale bar: 10 µm.



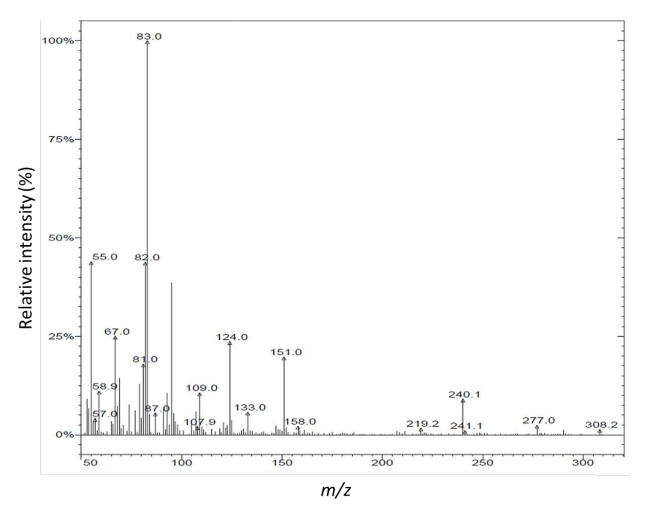
Supplementary Fig. S10. Amino acid sequence alignment of SmOPRs and other known OPRs.

Amino acid sequences were aligned using Clustal Omega. Identical and similar amino acids are highlighted in black and gray, respectively. The two specificity-determining residues are marked by red arrows. The aligned sequences include OPR3-like enzymes (AtOPR3: *Arabidopsis thaliana* OPR3, AEC06000; HbOPR: *Hevea brasiliensis* OPR, AAY27752; OsOPR7: *Oryza sativa* OPR7, Q6Z965; SIOPR3: *Solanum lycopersacum* OPR3, NP\_001233873; ZmOPR7: *Zea mays* OPR7, NP\_001105910; ZmOPR8, NP\_001105833), OPR1-like enzymes (AtOPR1: *Arabidopsis thaliana* OPR1, AEE35875; OsOPR1: *Oryza sativa* OPR1, Q84QK0; SIOPR1: *Solanum lycopersacum* OPR1, NP\_001234781), and SmOPRs. The numbering of SmOPRs was previously described by Li *et al.* (2009).

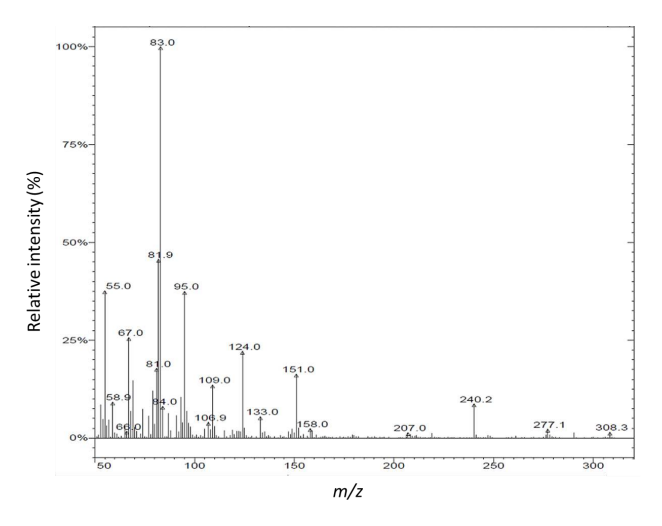


Supplementary Fig. S11. Phylogenetic tree of SmOPRs and previously reported OPRs.

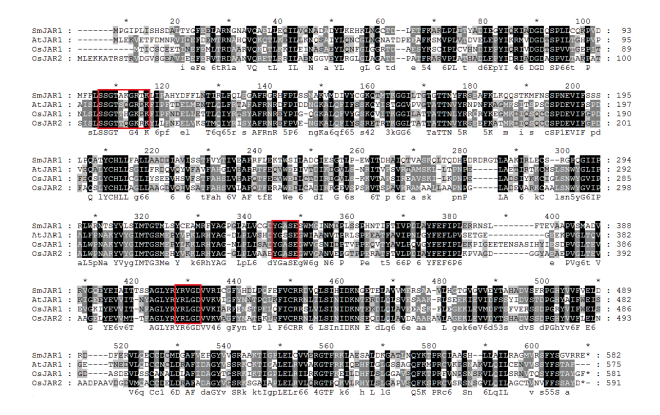
The OPR amino acid sequences were compared using Clustal Omega. The phylogenetic tree was visualized from the resulting alignment using TreeView X program. The analysis was performed with the following: OPR3-like enzymes (AtOPR3: *Arabidopsis thaliana* OPR3, AEC06000; HbOPR: *Hevea brasiliensis* OPR, AAY27752; OsOPR7: *Oryza sativa* OPR7, Q6Z965; SIOPR3: *Solanum lycopersacum* OPR3, NP\_001233873; ZmOPR7: *Zea mays* OPR7, NP\_001105910; ZmOPR8, NP\_001105833), OPR1-like enzymes (AtOPR1: *Arabidopsis thaliana* OPR1, AEE35875; OsOPR1: *Oryza sativa* OPR1, Q84QK0; SIOPR1: *Solanum lycopersacum* OPR1, NP\_001234781), and SmOPRs. The numbering of SmOPRs was previously described by Li et al. (2009).



Supplementary Fig. S12. GC-MS spectrum of the peak 1 of Fig. 6.

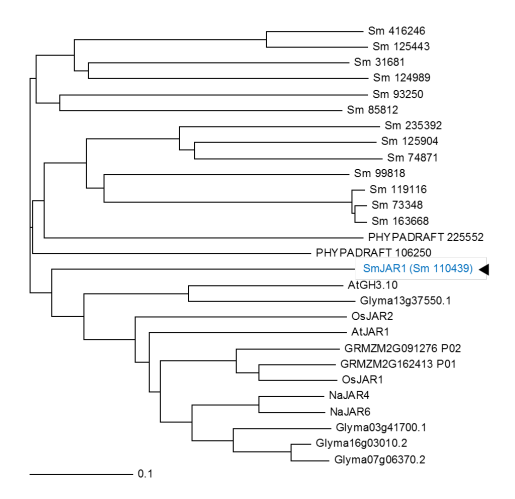


Supplementary Fig. S13. GC-MS spectrum of the standard methyl ester of OPC-8:0.



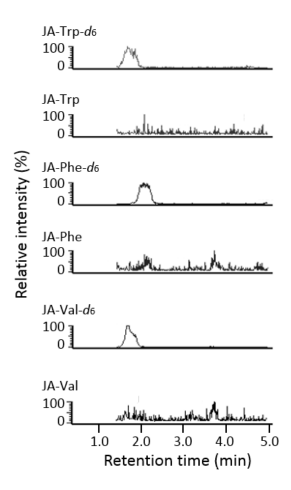
Supplementary Fig. S14. Amino acid sequence alignment of SmJAR1 with three GH3 family members.

Amino acid sequences were aligned using Clustal Omega. Identical and similar amino acids are highlighted in black and gray, respectively. The boxes indicates the three important motifs for enzymatic activity of GH3 proteins. The aligned sequences include AtJAR1 (*Arabidopsis thaliana*, AEC10684), OsJAR1 (*Oryza sativa*, LOC\_Os05g50890.1) and OsJAR2 (*Oryza sativa*, LOC\_Os01g12160.1).



Supplementary Fig. S15. Phylogenetic tree of JAR1-like proteins in S. moellendorffii with other JAR1s.

The tree was generated from amino acid alignment and constructed using Clustal Omega before visualizing with TreeView X program. The analysis was performed with the following: AtJAR1 (*Arabidopsis thaliana*, AEC10684), AtGH3.10 (*Arabidopsis thaliana*, At4g03400), OsJAR1 (*Oryza sativa*, LOC\_Os05g50890.1), OsJAR2 (*Oryza sativa*, LOC\_Os01g12160.1), Glyma16g03010.2 (*Glycine max*), Glyma07g06370.2 (*Glycine max*), Glyma03g41700.1 (*Glycine max*), Glyma13g37550.1 (*Glycine max*), NaJAR4 (*Nicotiana attenuate*, ABC87760.1), NaJAR6 (*Nicotiana attenuate*, ABC87761.1), GRMZM2G091276\_P02 (*Zea mays*), GRMZM2G162413\_P01 (*Zea mays*), PHYPADRAFT\_106250 (*Phsycomitrella patens*), and PHYPADRAFT\_225552 (*Phsycomitrella patens*).



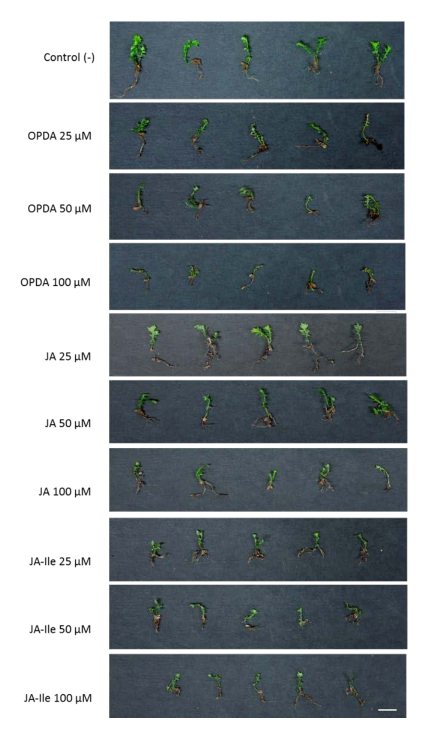
Supplementary Fig. S16. UPLC-MS/MS analysis of conjugates of JA and amino acid (JA-Trp, JA-Phe, and JA-Val) in the culture supernatant of *E. coli* expressing *SmJAR1*.

No obvious peak corresponding to JA-Trp, JA-Phe, and JA-Val was detected in the culture supernatant of *E. coli* expressing *SmJAR1*. Detailed experimental conditions are described in Materials and Methods.



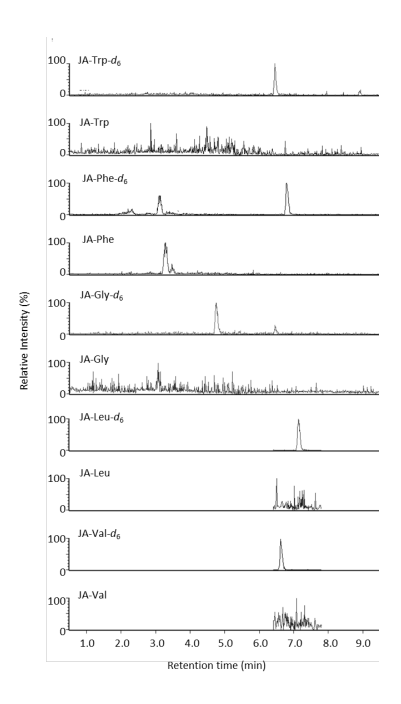
Supplementary Fig. S17. The morphology of a microphyll of *S. moellendorffii*.

The arrow indicates a bulbil.



Supplementary Fig. S18. Growth inhibitory activities of OPDA, JA, and JA-Ile in S. moellendorffii.

S. moellendorffii was treated with OPDA, JA, or JA-Ile at a concentrations of 25, 50, and 100  $\mu$ M. Detailed experimental conditions were described in Materials and Methods. Scale bar: 5 mm.



Supplementary Fig. S19. UPLC-MS/MS analysis of endogenous JA conjugation with other amino acids (Trp, Phe, Gly, Leu, and Val) in wounded *S. moellendorffii*.

No obvious peak corresponding to JA-Trp, JA-Phe, JA-Gly, JA-Leu, and JA-Val was detected in the extract of wounded *S. moellendorffii*.